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ABSTRACT

This study investigated the incorporation of DOM from seawater into <2 day-old sea ice in tanks filled with seawater alone or amended with DOM extracted from the microalga, *Chlorella vulgaris*. Optical properties, including chromophoric DOM (CDOM) absorption and fluorescence, as well as concentrations of dissolved organic carbon (DOC), dissolved organic nitrogen (DON), dissolved carbohydrates (dCHOs) and dissolved uronic acids (dUA) were measured. Enrichment factors (EFs), calculated from salinity-normalized concentrations of DOM in bulk ice, brine and frost flowers relative to under-ice water, were generally >1. The enrichment factors varied for different DOM fractions: EFs were the lowest for humic-like DOM (1.0–1.39) and highest for amino acid-like DOM (1.10–3.94). Enrichment was generally highest in frost flowers with there being less enrichment in bulk ice and brine. Size exclusion chromatography indicated that there was a shift towards smaller molecules in the molecular size distribution of DOM in the samples collected from newly formed ice compared to seawater. Spectral slope coefficients did not reveal any consistent differences between seawater and ice samples. We conclude that DOM is incorporated to sea ice relatively more than inorganic solutes during initial formation of sea ice and the degree of the enrichment depends on the chemical composition of DOM.

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Abstract

This study investigated the incorporation of DOM from seawater into <2 day-old sea ice in tanks filled with seawater alone or amended with DOM extracted from the microalga, *Chlorella vulgaris*. Optical properties, including chromophoric DOM (CDOM) absorption and fluorescence, as well as concentrations of dissolved organic carbon (DOC), dissolved organic nitrogen (DON), dissolved carbohydrates (dCHOs) and dissolved uronic acids (dUA) were measured. Enrichment factors (EFs), calculated from salinity-normalized concentrations of DOM in bulk ice, brine and frost flowers relative to under-ice water, were generally >1. The enrichment factors varied for different DOM fractions: EFs were the lowest for humic-like DOM (1.0- 1.39) and highest for amino acid-like DOM (1.10- 3.94). Enrichment was generally highest in frost flowers with there being less enrichment in bulk ice and brine. Size exclusion chromatography indicated that there was a shift towards smaller molecules in the molecular size distribution of DOM in the samples collected from newly formed ice compared to seawater. Spectral slope coefficients did not reveal any consistent differences between seawater and ice samples. We conclude that DOM is incorporated to sea ice relatively more than inorganic solutes during initial formation of sea ice and the degree of the enrichment depends on the chemical composition of DOM.

Introduction

When seawater freezes and sea ice forms, dissolved organic and inorganic matter is excluded from the crystalline ice structure and concentrates in the liquid brine phase. Brine may then drain into the water below, be involved in the formation of frost flowers on top of the ice, or remain trapped in brine channels and pockets within the sea ice (Petrich and Eicken, 2010). The quantity and chemical characteristics of brine affects both the spatial and temporal variabilities in the physico-chemical characteristics of sea ice, as well as the habitats for biological assemblages within ice (Thomas and Dieckmann, 2002). Dissolved organic matter (DOM) in brine is a major source of energy and nutrients for heterotrophic organisms within ice, and also significantly influences the optical properties of sea ice (Thomas and Dieckmann, 2010 and citations therein). When chromophoric DOM (CDOM) in brine absorbs ultraviolet radiation (UVR) and photosynthetically active solar radiation, it influences the energy budget of ice, the light availability and UVR exposure of organisms within and below the ice (Uusikivi et al., 2010).

In sea ice brines the concentration of DOM varies depending on brine volume (Petrich and Eicken, 2010; Thomas and Dieckmann, 2010): decreasing temperatures reduce the volume of brine and in turn increase the salinity and the concentrations of DOM in the brines above those in seawater. In contrast, when bulk ice, consisting of pure ice crystals and brine is melted, the concentrations of solutes (e.g., salinity and DOM) are lower in melted bulk ice than in seawater, because brine makes up only a small fraction of the total volume of ice. In order to assess the changes in the quantity of DOM in sea ice, without the effect of varying temperatures or sampling methods (brine vs. melted ice), the concentration of DOM in sea ice is frequently normalized to salinity (Giannelli et al., 2001; Granskog et al., 2004; Patsayeva et al., 2004; Müller et al., 2011). This has revealed that the DOM content of sea ice may include DOM from algal and bacterial production in sea ice (autochthonous; Underwood et al., 2010; Aslam et al., 2012a) in addition to that what is trapped in ice during its formation (allochthonous, Stedmon et al. 2007, 2011).

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77 Previously studies have suggested that DOM does not necessarily behave conservatively during
78 formation of sea ice compared to other dissolved constituents such as inorganic ions. As a
79 consequence the concentration of salinity normalized DOM can be higher (or enriched) in sea ice
80 compared with the under-ice water (Giannelli et al., 2001; Granskog et al., 2004; Patsayeva et al.,
81 2004; Müller et al., 2011). Despite these findings relatively little is known about the mechanisms of
82 enrichment during ice formation. One explanation is that the enrichment of DOM depends on
83 different diffusion rates of solutes to brine leaving sea ice through gravity drainage (Gross et al., 1987;
84 Reeburgh and Springer-Young, 1983). Such gravity drainage of brine may preferentially remove most
85 small molecules, such as inorganic ions, which diffuse at faster rates than DOM molecules with larger
86 size and lower diffusion rates (Granskog et al., 2004; Vancoppenolle et al., 2010; Maus et al., 2011).
87 Another explanation suggests that bacterial and algal extracellular polymeric substances (EPSs) form
88 gel-like structures that selectively retain DOM in sea ice (Raymond et al., 2007; Underwood et al.
89 2010; Krembs et al., 2011). Therefore, experimental evidence is needed to resolve the process of the
90 enrichment of DOM during ice formation, especially considering the enormous volume of sea ice that
91 forms, consolidates and subsequently melts in polar and sub-polar oceans and seas.

92

93 The present study addresses a potential enrichment of DOM during the first 48 h of ice growth in 1.2
94 m³ experimental mesocosms placed in a large environmentally controlled ice tank facility (at the
95 Hamburg Ship Model Basin HSVA: <http://www.hsva.de>). Absorption and fluorescence of CDOM, the
96 concentration of dissolved organic carbon (DOC), dissolved organic nitrogen (DON) and dissolved
97 carbohydrates (dCHOs) and uronic acids (dUAs) were analyzed from seawater, melted bulk ice, brine
98 and frost flowers to quantify different DOM fractions. Changes in the optical quality and the
99 molecular size of DOM during the formation of ice were investigated using analyses of the spectral
100 slope coefficients and size-exclusion chromatography of the CDOM. The seawater used was collected
101 from the North Sea, but in order to change both the quantitative and qualitative nature of the DOM,
102 high concentrations of DOM derived from disrupted algal cells were added to half of the mesocosms.

This enabled us to address the question of whether or not the quality of DOM affects the incorporation into sea ice. Aslam et al. (2012b) used the same experiment to investigate the effect of algal- DOM addition on bacterial communities and the production and dynamics of EPS. They described an incorporation of DOC and POC from the under-ice water into sea ice and the following change in the bacterial growth. The focus of the present study is to evaluate the abiotic incorporation of DOM from seawater to sea ice during the initial stages of ice growth where there was a minimal chance for the biological transformation of DOM (Aslam et al., 2012b). We describe a novel combination of methods that describe quantitative and qualitative changes of DOM during the initial ice formation. Understanding physico-biogeochemical processes during the ice formation are necessary to evaluate changes in the biomass, species composition and biochemistry of sea ice and help to estimate future conditions.

Materials and Methods

Experimental design and sampling routine

North Sea water was sampled by ship near Helgoland (54° 11' N, 7° 55' E) and transported in a cleaned (food-quality) road tanker to the HSVA test basin within 24 h, at a water temperature of $13 \pm 1^\circ\text{C}$.

Algal organic matter was produced by melting a frozen paste of freshwater algae *Chlorella vulgaris* (Varicon Aqua Ltd, U.K.) in artificial seawater with a salinity of 34. The suspension was sonicated in 500 ml batches at a sample temperature of 0°C using a Branson 450 Digital Sonifier (Branson Ultrasonics Corporation, Danbury, CT, USA). All suspensions were pooled and centrifuged at 12,235 g in a Beckmann Coulter centrifuge. The supernatant (algal-DOM) was collected and kept frozen (-18°C) until use. In this stock solution, the concentration of algal-derived DOC was 550 mmol L^{-1} .

The experimental setup consisted of 18 polyethylene (PE) bags supported by floating frames in an environmental test basin (Fig. 1). Each mesocosm was filled with 1.2 m^3 of unfiltered North Sea water

on September 30th 2009. 900ml of the algal-DOM of the stock solution was added to nine of the mesocosms (hereafter called SW+A), while the remaining mesocosms with just North Sea water are hereafter referred to as SW. All mesocosms were left to cool and mix (each mesocosm had a simple submerged pump installed), and initial samples were collected on the first sampling day (d0) on the 2nd of October after more than 24h of mixing. Freezing was initiated on d3 by spraying a fine mist of Milli-Q water (<1 L over whole environmental test basin) over the water surface (Giannelli et al., 2001) after the air temperature had been lowered to -13 °C, which was subsequently maintained for the rest of the experiment (± 2 °C). A polyvinyl chloride (PVC) tube was installed in each bag which was cleared of ice each day to maintain pressure equilibrium and ensure that the water was always in contact with the bottom of the ice, as well as to provide a convenient portal through which to sample under-ice water.

Water samples were collected from all mesocosms on d3, and thereafter water, ice and brine samples were collected from randomly chosen mesocosms on d4 and d5 with 1 to 3 replicates per treatment and sampling day. Each mesocosm was only sampled for ice and brine on one occasion, as the sampling compromised the integrity of the ice making the sampled mesocosm redundant (Fig. 1). Frost flowers were collected on d5. For the bulk ice samples, ice blocks were sawed from the ice sheet, floated carefully away from the remaining ice and sectioned within a minute into 2-3 layers depending on the ice thickness. This technique was employed to minimize the losses of brine that are characteristic of normal ice coring. The saw was cleaned before each use by sawing through other ice from the same depth horizon. The ice was melted at room temperature within 12 h in acid-washed PE buckets with the temperature of melt water never reaching above 0°C. Since the present study focused on the whole ice layer in comparison to under-ice water, we calculated a mean of measurements from all vertical ice layers in each mesocosm.

For brine sampling, sack holes (described by Papadimitriou et al., 2007) were drilled to a 6 cm depth with a Cherepanov ice ring of 20 cm diameter. Within 30 min after the drilling, brine was collected

from each sackhole using cleaned Teflon tubing and 20 ml syringes (without rubber parts) and these samples were processed in <4 h. Frost flowers were collected using the rim of an acid-washed PE container for scraping them from the ice surface. The frost flowers were melted at room temperature within 1 h, with the melt water never rising above 0°C, and again these samples processed in < 4h.

Sample treatment and analyses

Temperatures of water and brine were measured with a recently calibrated Testo® 110 thermometer. Temperatures ranged from 0.6°C on d0 to -1.9°C on d5 in water and from -3.1°C to -6°C in the brines. Optical properties of DOM were measured from 0.2 µm filtered (Millex-syringe filters; Millipore®) samples. Samples for liquid chromatography- size exclusion chromatography (LC-SEC) analysis were stored in pre-combusted glass scintillation vials in the dark at 4 °C. Samples for a DOC and DON were filtered through pre-combusted Whatman® GF/F filters and the filtrates were kept at -20°C until analyses. Salinity was measured using a SEMAT® Cond 315i/SET salinometer with a WTW Tetracon 325 probe at room temperature (Aslam et al., 2012b). DOC was measured by high temperature combustion on a MQ1000 TOC analyzer (Qian and Mopper, 1996), following the methods of Norman et al. (2011). DON was calculated by the subtraction of nitrate and ammonium from the total dissolved nitrogen (TDN). TDN was measured by standard colorimetric methodology (Grasshoff et al., 1983) on a LACHAT Instruments Quick- Chem 8000 autoanalyser (see Papadimitriou et al. (2007) for further details). Samples for dissolved carbohydrate (dCHO) analysis were desalted by dialysis (8 kDa membrane) against Milli-Q water (final salinity < 1), freeze dried and stored at -20 °C. Concentrations of dCHO were determined according to Underwood et al. (2010) and Aslam et al. (2012b). A standard carbazole assay was used to measure the concentration of dUA (Bellinger et al., 2005; Aslam et al., 2012b).

Absorption spectra of CDOM were measured in a 10 cm quartz cell over a 200 to 700 nm range in 1 nm increments and a slit width of 2 nm with a Shimadzu UV-2101 spectrophotometer. After the

samples had warmed to room temperature, three replicates of each sample were measured against ultrapure water (MilliQ, resistivity 18.2MΩ cm) blank. CDOM was described by the absorption coefficient, a (m^{-1}), at wavelength λ (nm) that was calculated using

$$a_{\text{CDOM},\lambda} = 2.303A_{\lambda} 0.1^{-1} \quad (1),$$

where A_{λ} is the mean absorbance at wavelength λ .

The spectral slope coefficient S (μm^{-1}) for the ranges 250-450 nm, 275-295 nm and 350-400 nm was calculated by using a nonlinear fit in Matlab based on the following equation (Stedmon et al., 2000):

$$a_{\text{CDOM},\lambda} = a_{\lambda,0} e^{S(\lambda,0 - \lambda)} \quad (2).$$

where $a_{\lambda,0}$ is the absorption coefficient at the wavelength $\lambda,0$ describing the shortest end of spectral range. The coefficients of determination for the fits were >0.99 . The slope ratio (S_R) was the ratio of $S_{275-295}$ to $S_{350-400}$ (Helms et al. 2008).

Fluorescence was measured in a 1 cm quartz cell using a Varian Cary Eclipse spectrofluorometer with an integration time of 0.1 s and a scan speed of 1200 nm min^{-1} . The voltages changed depending on the concentration of the sample between 850 and 1000V but were corrected later by calibration to the Raman scatter signal (Lawaetz and Stedmon, 2009). Excitation ranged from 240 to 450 nm in 5 nm increments and emission from 300 to 550 nm with 2 nm increments. The slit width was set to 5 nm for excitation and emission scans. Measurements and instrumental corrections were done according to Stedmon and Bro (2008). Excitation Emission matrices (EEMs) of all 116 samples were characterized by 6 fluorescent components using parallel factor analysis (PARAFAC) and the DOMFluor toolbox (Stedmon and Bro, 2008). The components were validated by split half analysis and random initialization. The fluorescence intensities in the subsequent data analysis refer to the maxima fluorescence signals of each component (Fmax).

The molecular size distribution of CDOM was analyzed by LC-SEC as described by Müller et al. (2011) using a TSK G3000SWxl column (7.8 mm \times 30 cm, 250 Å pore size, 5 μm particle size) and a

TSK guard column (7.5 mm × 7.5 mm), with a KH₂PO₄ (2.7 g L⁻¹) and Na₂HPO₄·2H₂O (3.56 g L⁻¹, pH 6.85) buffer. Measurements were performed using absorbance detection at 254 nm with a Hewlett Packard 1100 series high-performance liquid chromatography (HPLC) instrument. All samples were diluted with ion-exchanged Milli-Q water to a final salinity of 9 to avoid a shift in molecular size due to changes in salinity (Specht and Frimmel, 2000; Her et al., 2002). A baseline correction was performed by normalizing all chromatograms to the absorption at the retention time (Rt) of 12 min. Salinity-normalization was done by setting the salinity-dependent maximal value of each sample to 1 (Minor et al., 2002). Rts were 7.7 min for Blue Dextran 2000 (2×10⁶ g mol⁻¹), 16.3 min for tyrosine (181.2 g mol⁻¹), 16.5 min for phenylalanine (165.1 g mol⁻¹) and between 12 and 15.5 min for Nordic fulvic acid (Müller et al., 2011). We limited our interest in the range of Rt from 13.5 to 15.5 min representing the molecular size fraction of fulvic acids.

Salinity-normalized concentrations of DOM fractions collected from different compartments of ice or water ($X_{norm,i}$) were calculated by:

$$X_{norm,i} = \frac{X_i}{S_i} 33 \quad (3),$$

where X represents the DOM fraction (DOC, DON, dCHO, dUA, CDOM or maximum intensities of fluorescent components), the subscript i represents the sample type (water, bulk ice, brine or frost flowers) and 33 was the initial salinity of North Sea water.

The enrichment factor of DOM fractions, $EF(X,i)$, was calculated by:

$$EF(X,i) = \frac{X_{norm,i}}{X_{norm,water}} \quad (4),$$

where $X_{norm,i}$ and $X_{norm,water}$ are the salinity normalized concentrations of the DOM fractions in each sample type and in water below the ice, respectively. $EF(X)$ notation refers to DOM fractions specified by X without referring to any specific type of sample collected from ice. If $EF > 1$, a DOM fraction in ice is considered to be enriched and conversely a $EF < 1$ represents depletion relative to seawater.

The main aim in this study was to assess potential changes in DOM during its incorporation to newly formed sea ice. For this assessment we compared samples collected from sea ice (bulk ice, brine or frost flowers) to samples collected from water using t-tests for independent samples (SPSS; $p \leq 0.05$).

Results

Characterization of initial water

The addition of algal-DOM to North Sea water changed the absorption spectrum of CDOM, the excitation-emission matrix (EEM) of fluorescent DOM and the size distribution of humic-like dissolved organic matter (Fig. 2). The algal-DOM had little influence on CDOM absorption at wavelengths above 300 nm, except for a small peak at 410 nm (14 % increase), however, the absorption at 255nm doubled (Fig. 2a). The spectral slope coefficient $S_{250-450}$ was $18.1 \pm 0.3 \mu\text{m}^{-1}$ (\pm refers to standard error throughout the results) in the SW mesocosms, but higher ($26.6 \pm 0.1 \mu\text{m}^{-1}$) in the mesocosms with algal-DOM, because the lower spectral range of $S_{250-450}$ matched an absorption peak of algal-DOM (Fig. 2a). S_R values were 1.19 ± 0.04 in the SW treatment and 1.96 ± 0.04 in the SW+A-treatment. The addition of algal-DOM increased the intensity of fluorescence at shorter emission wavelengths (Fig. 2b). After the addition of algal-DOM, the size distribution chromatography showed an increase in the absorbance by 4% at the retention time of 14.6 (peak 1) and by 22% at the retention time of 15.2 min (peak 2; Fig. 2c). The molecular size distribution was therefore shifted towards smaller molecules in the SW+A treatment. The addition of algal-DOM increased the initial concentrations of DON from $3.9 \pm 0.2 \mu\text{mol L}^{-1}$ to $44.1 \pm 1 \mu\text{mol L}^{-1}$. The corresponding change for DOC was from $109.4 \pm 3.2 \mu\text{mol L}^{-1}$ to $380 \pm 5.1 \mu\text{mol L}^{-1}$. dCHO concentrations were $25.2 \pm 1.3 \mu\text{mol L}^{-1}$ and $31.7 \pm 1.1 \mu\text{mol L}^{-1}$ in the SW- and SW+A treatments, respectively, and the concentrations of dUA were $13.7 \pm 0.7 \mu\text{mol L}^{-1}$ and $17.6 \pm 1.4 \mu\text{mol L}^{-1}$ in the SW- and SW+A treatments, respectively.

The PARAFAC modeling of fluorescent DOM (FDOM) separated the pool of FDOM into 6 components (Fig. 3a): C2 (ex/em = 340/430 and 240/430), C3 (310/385 and 240/385) and C4 (245/500 and 380/500) were similar to humic-like components (Coble 1996; Murphy et al., 2008; Yamashita et al., 2010; Stedmon et al., 2011). The remaining components had characteristics similar to amino acids. C5 (ex/em=275/315) was similar to a tyrosine-like component (Murphy et al., 2008), while C1 (280/360) and C6 (240/350 and 290/350) were similar to tryptophan-like components which have excitation maxima below 240 nm and at 275 nm and an emission maximum at 350 nm (Stedmon and Markager, 2005; Murphy et al., 2008; Stedmon et al., 2011).

In the SW +A treatment waters, the intensity of fluorescence was highest for C1 (Fig. 3b). The PARAFAC analysis of FDOM indicated that the introduction of algal-DOM increased the concentrations of amino acid-like DOM up to 16-fold and humic-like DOM up to 1.3-fold (Fig. 3b).

Dynamics of salinity during the experiment

The experiment consisted of the cooling phase (d0 to d3) prior to the ice formation and the freezing phase with the first two days of ice (d4 and d5, indicated by the grey background in Fig. 4). The development of the ice thickness and the salinity was similar in both treatments (Fig. 4 inset). The salinity of the four different sample types (water, bulk ice, brine and frost flowers) showed large differences mainly caused by the exclusion of salts from bulk ice and their enrichment in brine (Fig. 4). In order to compare the behavior of DOM fractions (X) across the sample types with different salinity, their concentrations were normalized to the initial salinity of seawater which was 33 (Eq. 3)

Quantitative changes in DOM during freezing

DOC_{norm} and DON_{norm} decreased throughout the experiment in the SW+A but not in the SW-treatment and were typically significantly higher in bulk ice, brine and frost flowers than in water as indicated by the asterisks in Fig. 5. These results indicate that DOC_{norm} and DON_{norm} were generally enriched in

all sample types collected from sea ice relative to water. $EF(DOC)$ or $EF(DON)$ for the samples collected from ice (bulk ice, brine, frost flowers) ranged from 1.08 to 3.64 (Fig. 6a-d; Eq. 4).

After the formation of ice, $CDOM_{norm}$ was consistently higher in bulk ice and frost flowers than in water (Fig. 7a-b). $EF(CDOM)$ of all sample types ranged from 0.96 to 1.30 (Fig. 6 e-f).

$FDOM_{norm, bulk\ ice}$ exceeded $FDOM_{norm, water}$ for all 6 fluorescent components in the SW-treatment, but only for C4 and C6 in the SW+A-treatment (Fig. 8 left column and Fig. 8 h and l, respectively). In the SW+A-treatment, CI_{norm} of bulk ice was lower than that in SW indicating a depletion of C1 in bulk ice (Fig. 8b). The enrichment factors for the fluorescent components ranged from 0.83 to of 7.45 (Fig. 6 g-r).

After the ice formation, $dCHO_{norm}$ and dUA_{norm} were typically higher in the bulk ice than in water (Fig. 9 a-d). $dUA_{norm, brine}$ instead was lower $dUA_{norm, water}$ (Fig 9 c-d). The enrichment factors for dCHO and dUA ranged from 0.99 to 2.54 in bulk ice and from 0.5 to 1.01 in brine (Fig. 6 s-v).

Changes in spectral slope and molecular size distribution

Spectral slopes and slope ratios calculated for ice samples were generally similar to those measured in water (Fig. 7 c-j), however, in a few cases the slopes were significantly different between water and bulk ice, brine or frost flowers (Fig. 7 e-j). These differences were not consistent in both treatments suggesting that freezing did not change spectral slopes in a consistent manner (Fig. 7 e-j).

In the LC-SEC analysis, the variation in the UV absorbance at $R_t < 13.5$ min did not exceed the standard deviation of the replicated measurements (Fig. 2c). The later parts of the chromatograms ($R_t = 13.5$ to 15.5 min) varied most and therefore we examined R_t and absorbance values of P1 and P2 found in this region of the chromatogram in detail (Fig. 2c, Fig. 10). The intensities of P1 and P2 were

not significantly different between water and bulk ice, brine or frost flowers (Fig. 10 e-h). Rts of P1 and P2 were often longer in bulk ice, brine and frost flowers than in water (Fig. 10 a-d) indicating shift in the molecular mass distribution of DOM towards smaller molecules.

Enrichment factors

For further analysis of enrichment factors reported in Fig.6, the EFs for humic type DOM (CDOM, C2, C3 and C4) and amino acid-like fluorophores (C1, C5 and C6) were combined and examined with EFs for the other DOM fractions (Fig. 11). Among these DOM fractions, $EF(humics)$ ranged from 1.0 to 1.39 in different sample types. The highest EFs and highest variability among replicates were found for amino acid-like DOM (see $EF(aa)$ in Fig. 11). The $EF(dCHO)$ and $EF(dUA)$ ranged from 0.73 to 2.69 (Fig. 11). These results indicate that although all fractions of DOM were generally enriched in newly formed ice, the degree of enrichment differed between the various DOM fractions.

EFs were generally lower in the treatment with algal-DOM than in the treatment without it (Fig. 11; independent samples t-test; $p < 0.05$). Enrichment of DOM was generally lower in the brines compared with the bulk ice samples (Fig. 11; independent samples t-test; $p < 0.05$). In brines, the EFs were sometimes below 1.0 (e.g., $EF(dUA, \text{brine})=0.73$) indicating a depletion of DOM compared to water. EFs were higher in frost flowers (2.26 ± 0.26 , mean \pm SE for all DOM fractions) compared with bulk ice or brines (independent samples t-test; $p < 0.05$). These results indicate that EFs in frost flowers were up to 4 times higher than those in bulk ice samples and 7 times higher than in brines ($EF(C6)$, Fig.6r).

Discussion

Selective incorporation of DOM fractions during ice formation

The present study investigated the behavior of different DOM fractions in bulk ice, brine and frost flowers during initial ice formation. The results show that all investigated DOM fractions (DOC,

DON, CDOM, FDOM, dCHO, dUA) are generally enriched in bulk ice when normalized to the salinity of water. As in our study, enrichment of CDOM, DOC and DON has been observed in natural ice of unknown age in the Baltic Sea (Granskog et al., 2004; Stedmon et al., 2007) and the Southern Ocean (Meiners et al., 2009; Stedmon et al., 2011; Underwood et al., 2010; Norman et al., 2011). Since our results concern new ice (< 2 days old), they indicate that DOM is enriched already during the formation of ice mainly through abiotic physico-chemical processes.

Natural older sea ice can have $EF(DOC, \text{bulk ice})$ of approximately 6.71 ± 2.10 and $EF(DON, \text{bulk ice})$ as high as 11.58 ± 5.70 (calculated from Norman et al., 2011). The values in new ice measured experimentally in the present study are substantially lower ($EF(DOC, \text{bulk ice}) = 2.25 \pm 0.48$; $EF(DON, \text{bulk ice}) = 1.72 \pm 0.11$). This likely reflects the contribution of autochthonous production in natural ice (Fig. 11, Stedmon et al., 2007; Norman et al., 2011).

The enrichment of CDOM in this study ($EF(CDOM, \text{bulk ice}) = 1.12 \pm 0.03$) is similar to that in young ice formed in brackish Baltic Sea water ($EF(CDOM, \text{bulk ice}) = 1.34 \pm 0.16$; Müller et al., 2011). Similar to the patterns seen for CDOM, fluorescent components related to humic substances also have low EFs between 1.16 and 1.27 in young ice formed from water of the Baltic Sea and the North Sea (Müller et al., 2011, this study). In both studies the amino-acid-like components had the highest EFs among the FDOM-components examined: 1.53 in the Baltic Sea (Müller et al. (2011) and 2.72 ± 0.47 in the North Sea (present study). The mean of $EF(dCHO, \text{bulk ice})$ and $EF(dUA, \text{bulk ice})$ of 1.9 ± 0.191 in our study is similar to 1.61 ± 0.3 for exopolymers in newly formed artificial sea ice (Ewert and Deming, 2011). When our results are combined with earlier studies (Ewert and Deming, 2011, Müller et al., 2011), it seems that the degree of enrichment depends on the chemical characteristics of DOM with the lowest enrichment for humic type-DOM and highest enrichment for amino acid-like DOM.

Is there a mechanistic explanation for the enrichment of DOM in new sea ice?

374

375 This study revealed that while DOM is incorporated in newly formed sea ice, the size distribution of
376 humic-like DOM changes towards smaller molecules. We are not aware of other studies, which have
377 measured changes in the molecular size distribution of DOM during freezing. However, earlier studies
378 concerning the freeze fractionation of inorganic ions have suggested that freezing rejects preferably
379 smaller ions with larger diffusion coefficients (e.g., $2 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ for K^+) than larger ions with
380 smaller diffusion coefficients (e.g., $0.5 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ for SO_4^{2-} ; Granskog et al., 2004; Maus et al.,
381 2011). The diffusion coefficients of humic-like DOM, shown in our analyses, can range over four-fold
382 like those of major inorganic ions in seawater (Hassellöv, 2005; Siripinyanond et al. 2005). If
383 diffusion plays a major role in the process of selective rejection of DOM as suggested for inorganic
384 ions, we should have observed a shift towards larger molecules in the samples collected from ice in
385 our LC-SEC measurements. On the contrary, our study indicates a shift towards smaller molecules in
386 the molecular size distribution of DOM during incorporation to sea ice. This indicates that diffusion
387 cannot be the major factor that controls the selective rejection of humic-like DOM in ice. Additionally,
388 according to Aslam et al. (2012b) different size-classes of dCHOs enrich similarly to the new sea ice.
389 Hence, this study and that of Aslam et al. (2012b) indicate that freezing changes the molecular size of
390 distribution of DOM only little when DOM from seawater is incorporated in newly formed sea ice.

391

392 Both dUAs and dCHO examined in this study are part of the pool of material that also included
393 dissolved EPS and transparent exopolymeric particles (TEP) and can result in the coagulation of
394 DOM and potentially enhance the enrichment of DOM in sea ice (Chin et al. 1998; Engel et al., 2004;
395 Verdugo et al. 2004). In our experiment, the addition of algal-DOM doubled the concentration of
396 dUA and dCHO in the SW+A treatment compared to the SW-treatment. The enrichment of DOM
397 would have been expected to be higher in the algal-DOM treatment if dUA and dCHO enhance
398 enrichment of DOM (Aslam et al. 2012b). Contrary to this expectation, EFs were lower in the SW+A
399 than in the SW-treatment. Our observation suggests that dUA and dCHO did not influence enrichment

of DOM during the initial formation of ice, but that does not exclude such possibility in older sea ice, where algal derived DOM and EPS would play more of a role than in these short term experiments.

In this study, the salinity normalized concentrations of DOM, and the EFs calculated from them, were lower in brine than in bulk ice, as also observed for natural sea ice in the Southern Ocean (Underwood et al., 2010; Norman et al. 2011). All three studies collected brine with the sackhole-technique (Papadimitriou et al. 2007). When a sackhole is filled with brine having an $EF(DOM, \text{brine})$ lower than in the original bulk ice, the $EF(DOM, \text{bulk ice})$ in the surrounding bulk ice must increase simultaneously in less than 30 minutes, the time used for collection of brine in this study. Differences in the diffusion rates among solutes may not be able to explain such rapid changes in $EF(DOM)$ during the filling of a sackhole. One alternative potential mechanism is that there was selective drainage: i.e. brine from large brine channels, but with low $EF(DOM)$, drain more effectively into a sackhole than disconnected or very small brine channels with higher $EF(DOM)$. The change in $EF(DOM, \text{brine})$ may be also explained by a selective drainage of the most soluble solutes with low $EF(DOM)$ moving with a hydraulic flow to a sackhole. In this case, the less soluble solutes, or their insoluble forms in highly saline brine, will result in high $EF(DOM, \text{bulk ice})$. A poor solubility of DOM at high salinities may be explained e.g., by the association between multivalent cations and negatively-charged functional groups of DOM (Chave and Suess, 1970). In the latter case, both salts and DOM are converted to insoluble forms, which are likely overlooked when examining filtered samples alone (e.g., in the present study). If a similar drainage of brine with low $EF(DOM, \text{brine})$ into the under-ice water takes place during initial ice formation, it can result in the observed enrichment of the remaining DOM in new ice (Giannelli et al., 2001; Müller et al., 2011; this study).

Frost flowers are initially formed from water evaporating from brine into the colder atmosphere and only then, brine is drawn up onto the frost flower crystals by capillary actions (Perovich and Richter-Menge, 1994; Martin et al., 1996; Rankin et al., 2002). Therefore, depending on the temperature, the salinity of frost flowers can be as low as the seawater salinity in the early stage of frost flower

formation, as found in the present experiment. Despite the low salinity in frost flowers, there was a significant enrichment of DOM compared to under-ice water and brine from North Sea water (Fig. 6, Fig. 11, Bowman and Deming, 2010; Aslam et al., 2012b). Hence, we suggest that there is a second selective rejection process that occurs during the rapid freezing of brine on the ice surface, similar to the process during sea ice formation.

In this study various components of DOM are enriched in ice relative to water, even in the treatment with algal-DOM, which increased the concentration of DOM (DON and amino acids in particular) in water. The introduction of such potentially biologically labile DOM increased microbial activity in water and sea ice of the SW+A treatment (Aslam et al. 2012b) resulting in a decrease of DOC and DON in water later in the experiment than the initial stages reported here. It is possible, that the observed changes in the EFs of labile DOM components and in the molecular mass distribution and $S_{250-450}$ during sea ice formation are at least partly related to elevated microbial activity in SW+A – treatment.

Conclusions

This study shows that there is a quantitative enrichment for the whole DOM pool regardless of its initial concentration during initial freezing of seawater, which is probably based on a physico-chemical process. The enrichment factors vary between 1.0 and 1.39 for humic-like material, but are higher and more variable for amino-acid like DOM fractions. The results indicate that diffusion has only a minor effect on the enrichment process and also the concentration of carbohydrates in the seawater did not explain the enrichment behavior in this short-term experiment. Enrichment of DOM in melted ice differs from that in brine suggesting that the DOM concentration and composition varies among well- connected brine channels and the smallest brine channels and inclusions. These differences in the selective rejection of DOM from bulk ice and brine also mean that the method of sackhole sampling does not necessarily succeed in extracting brine that is representative of *in situ* conditions. Frost flowers had a similar DOM composition to that of the brines, but at higher

concentration, relative to salt, suggesting that a second fractionation process occurs during frost
flower formation.

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 596



597

598 Fig. 1: The experimental mesocosms floating in an environmentally controlled ice tank of the
599 Hamburg Ship Model Basin. The foam pads on some of the mesocosms mark the compromised ones
600 after they have been sampled. Note that the lights were on only during sampling (at maximum 2 hours
601 per day) and the rest of the time, the tanks were in darkness.

602

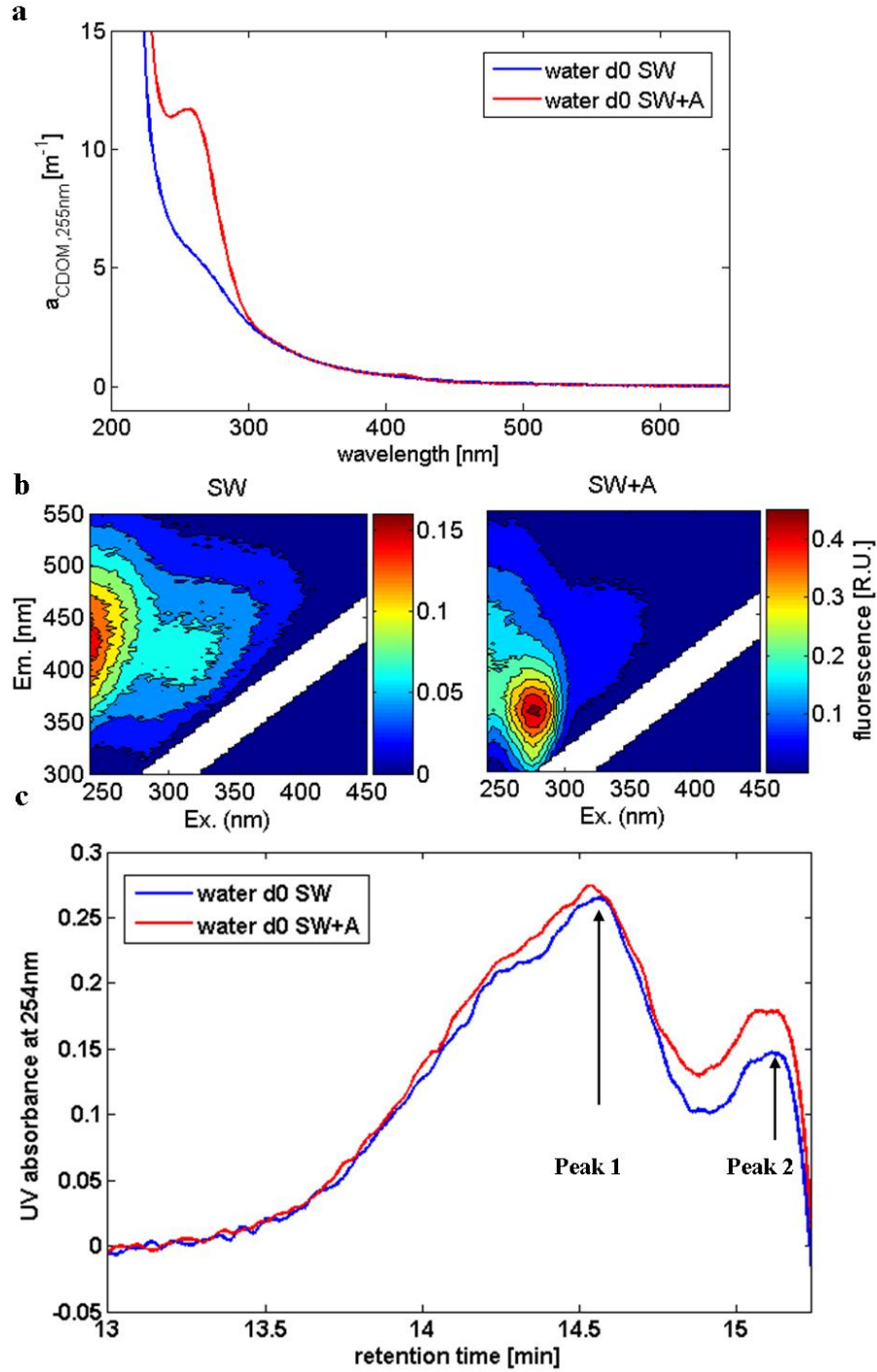


Fig. 2: The initial CDOM absorption spectra (a), fluorescence excitation emission matrices (b) and LC-SEC chromatograms (c) in North Sea water alone (SW) and with algal-DOM added (SW+A). The curves for CDOM (a) and LC-SEC (c) show means ($n = 7 - 8$). The standard errors (SE) of means were less than the thickness of curves.

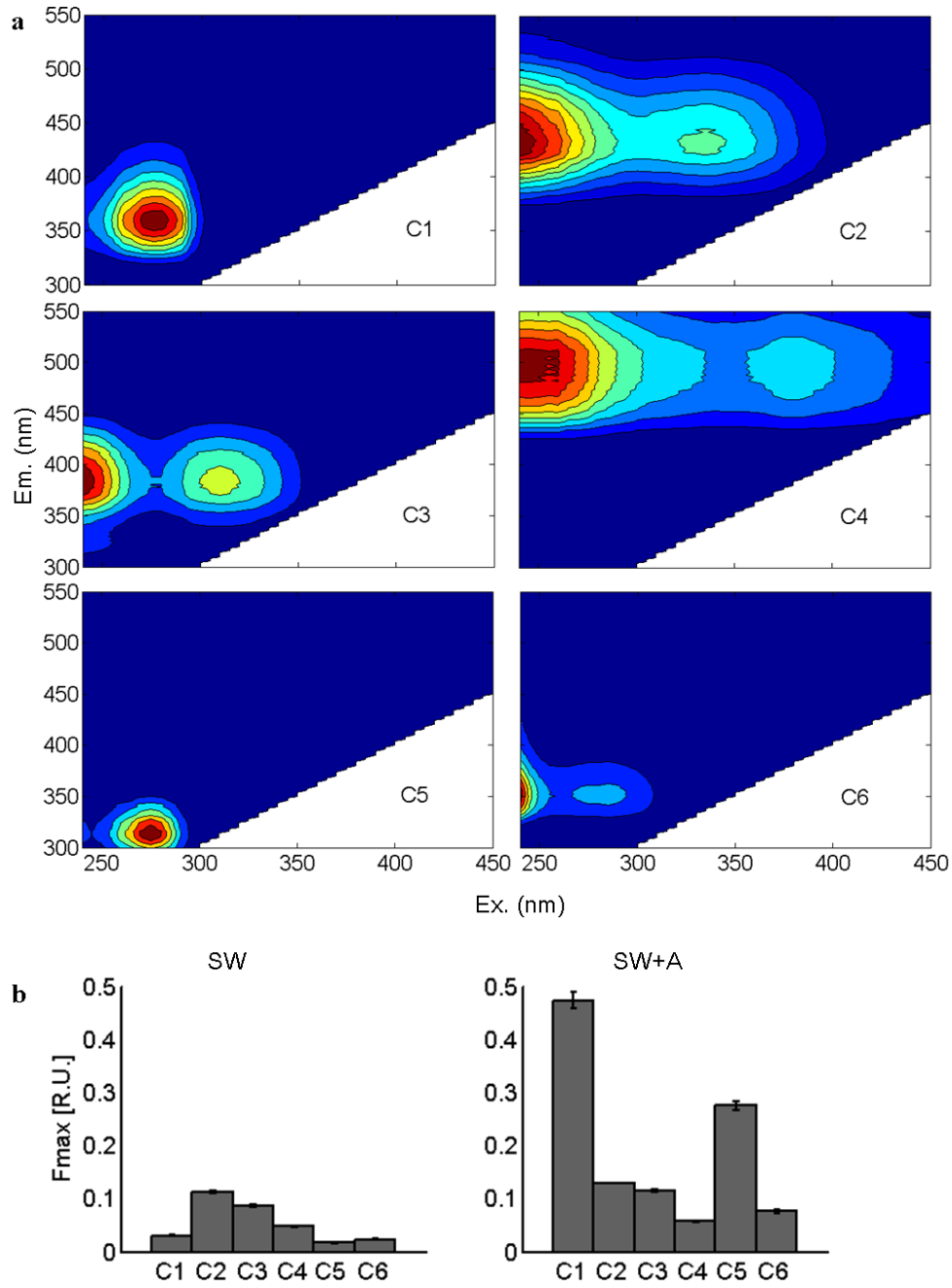


Fig. 3: The six fluorescent components identified from the pool of 116 samples (a) and their fluorescence intensities in the beginning (d0) of SW and the SW+A treatments (b). Fmax refers to mean \pm SE of maximum fluorescent intensities of each component in the replicated mesocosms ($n = 7 - 8$).

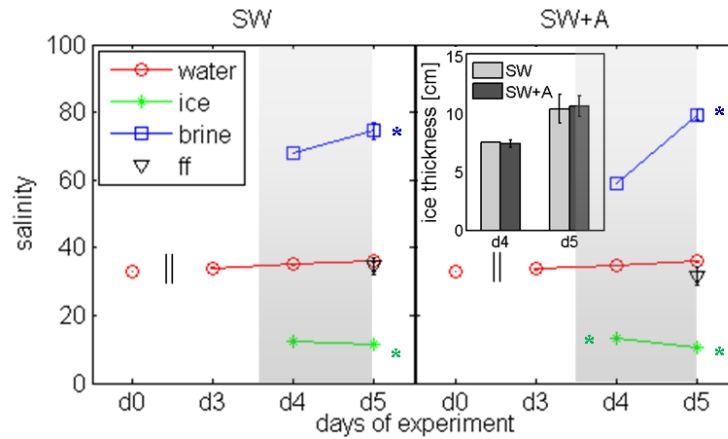


Fig. 4: The salinity in water, brine, bulk ice and frostflowers (ff) over the time of the experiment for each treatment together with the ice thickness on d4 and d5. Error bars show the standard error (SE) among replicated mesocosms ($n = 1 - 8$). Significant differences in salinity (t-test for $n > 1$, $p < 0.05$) of brine, bulk ice or frost flowers compared to water are indicated by *. The grey shading indicates the freezing phase of the experiment. The treatments with North Sea water alone and with algal-DOM are labelled SW and SW+A, respectively. The inset shows the mean ice thickness of SW and SW+A at d4 and d5 with the SE represented by error bars.

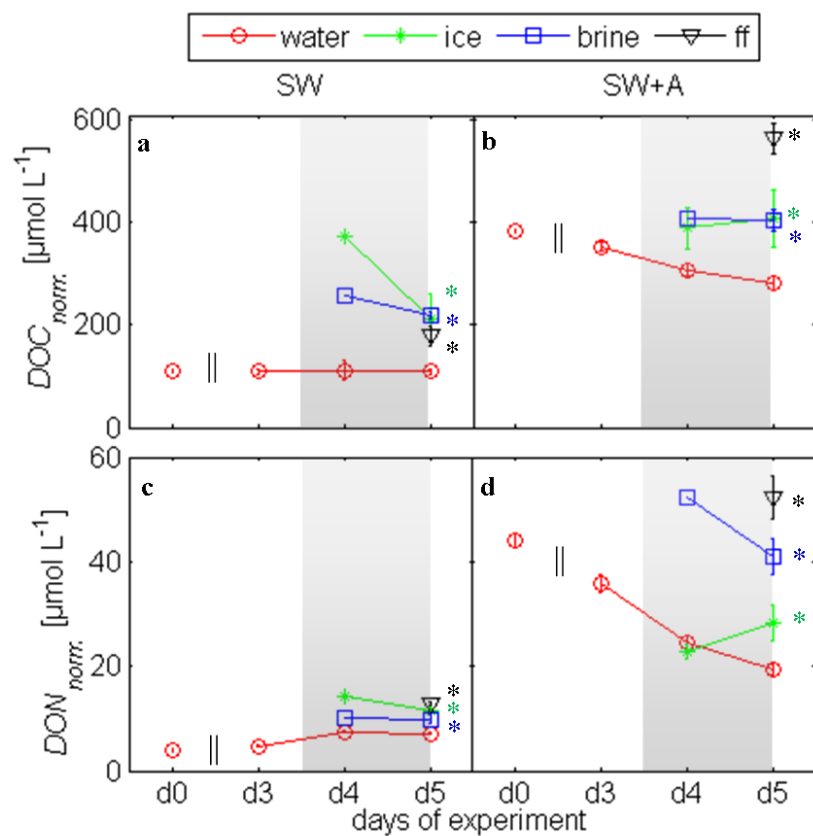


Fig. 5: Temporal development of DOC and DON after salinity normalization to 33. See Fig. 4 for more detailed explanation.

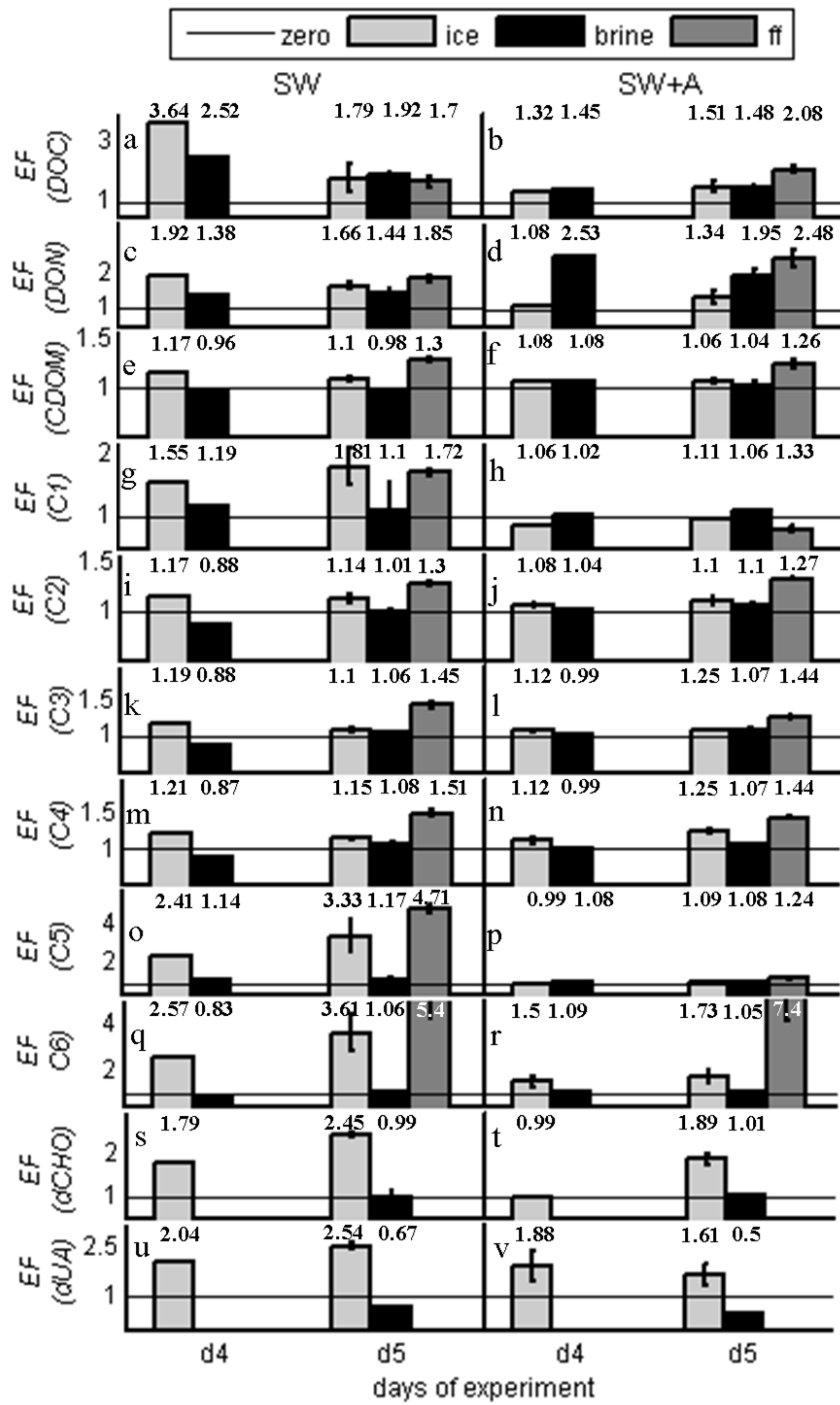


Fig. 6: Enrichment factor (EF) of DOC, DON, $a_{CDOM,255nm}$, fluorescent components C1 to C6, dCHO and dUA in bulk ice (light grey), brine (black), and frost flowers (dark grey) on d4 and d5 (see Eq. 4). The black line visualizes the point of zero enrichment. Error bars show the standard error among replicated mesocosms (n = 1 - 3). Numbers on top of each bar show the mean EF. Please note the different scales.

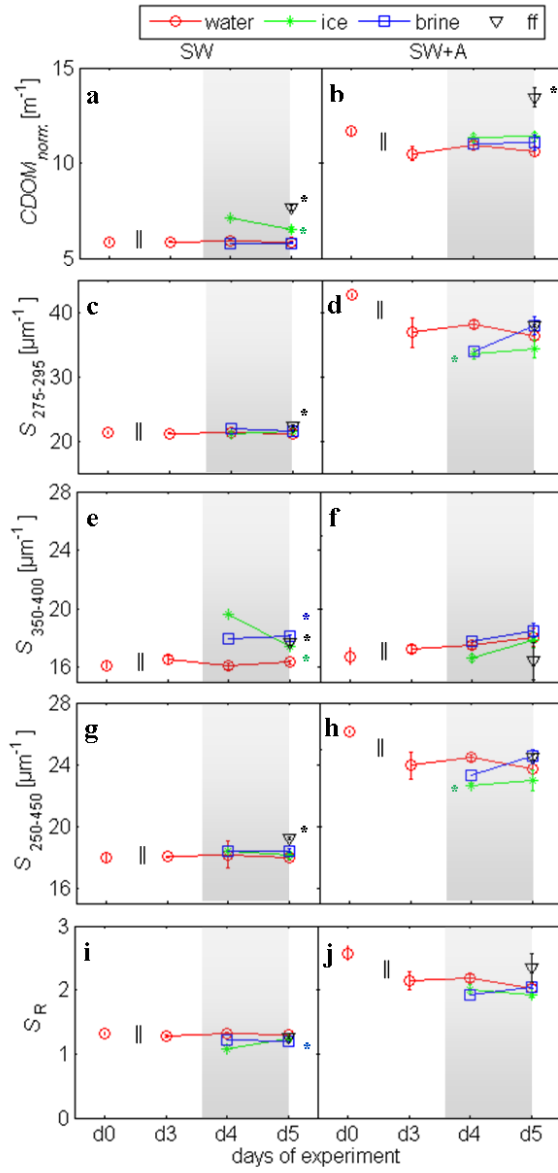


Fig. 7: Temporal development of absorption coefficient of CDOM at 255nm normalized to water salinity, $CDOM_{norm}$ (a-b), the spectral slope coefficient $S_{275-295}$ (c-d), $S_{350-400}$ (e-f), $S_{250-450}$ (g-h) and S_R values (i-j). The notation used and the salinity normalization are explained in Fig.4 and Eq. 3, respectively.

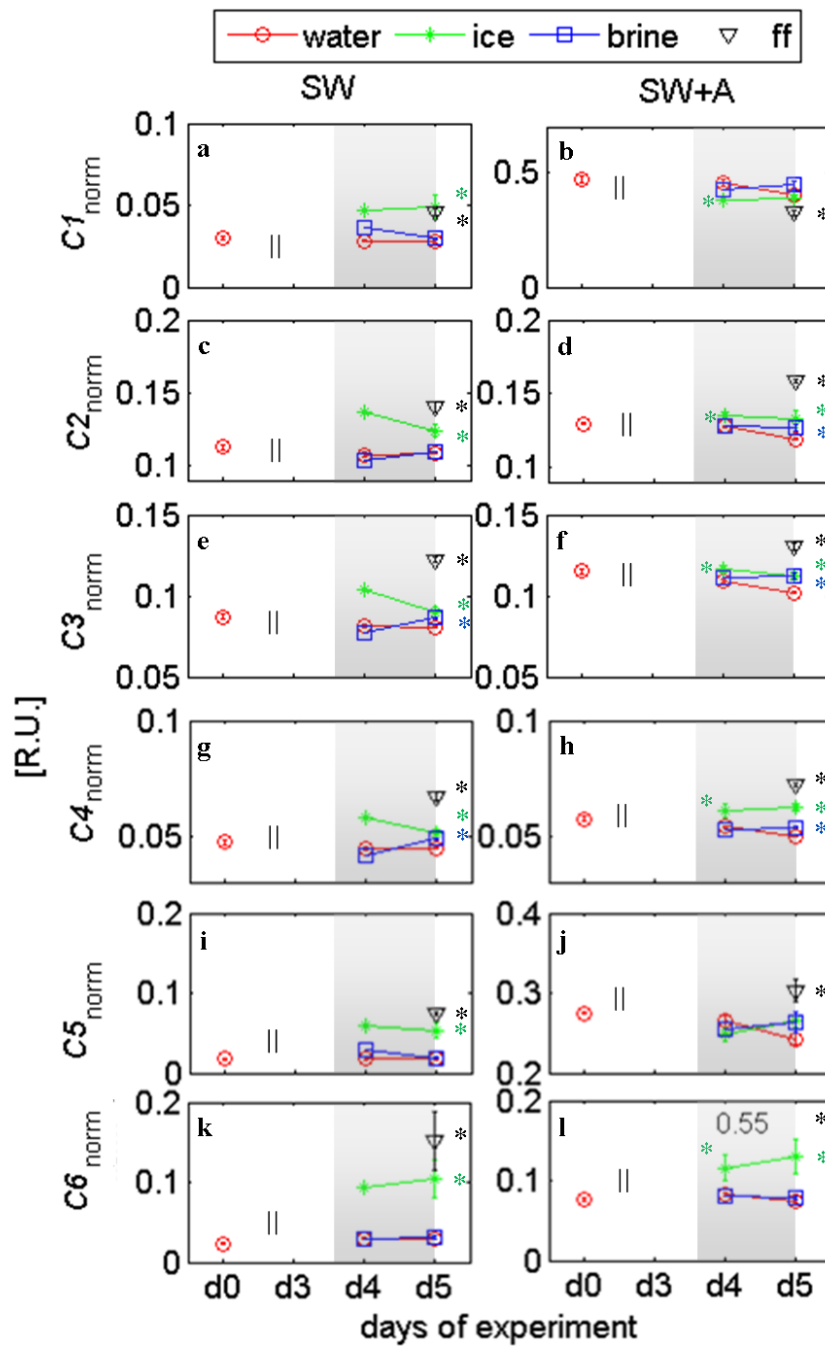


Fig. 8: Temporal development in the intensity of maximum fluorescence of the components 1 to 6 after salinity normalization to 33. Note the different scales between the treatments for C1 and C5 as well as the off-scale value 0.55 for frost flowers in panel l. See Fig. 4 for more detailed explanation.

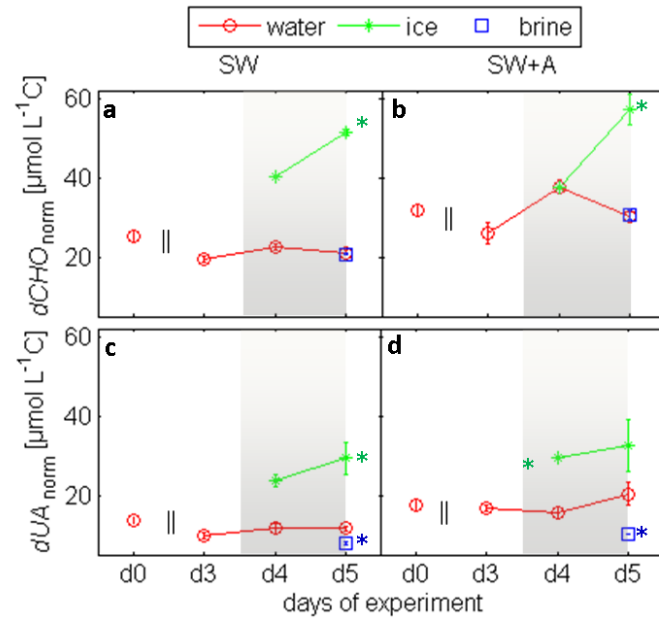


Fig. 9: Temporal development of salinity-normalized concentrations in dissolved carbohydrates (Fig.8 a-b) and dissolved uronic acids (Fig.8 c-d). See Fig. 4 for more detailed explanation.

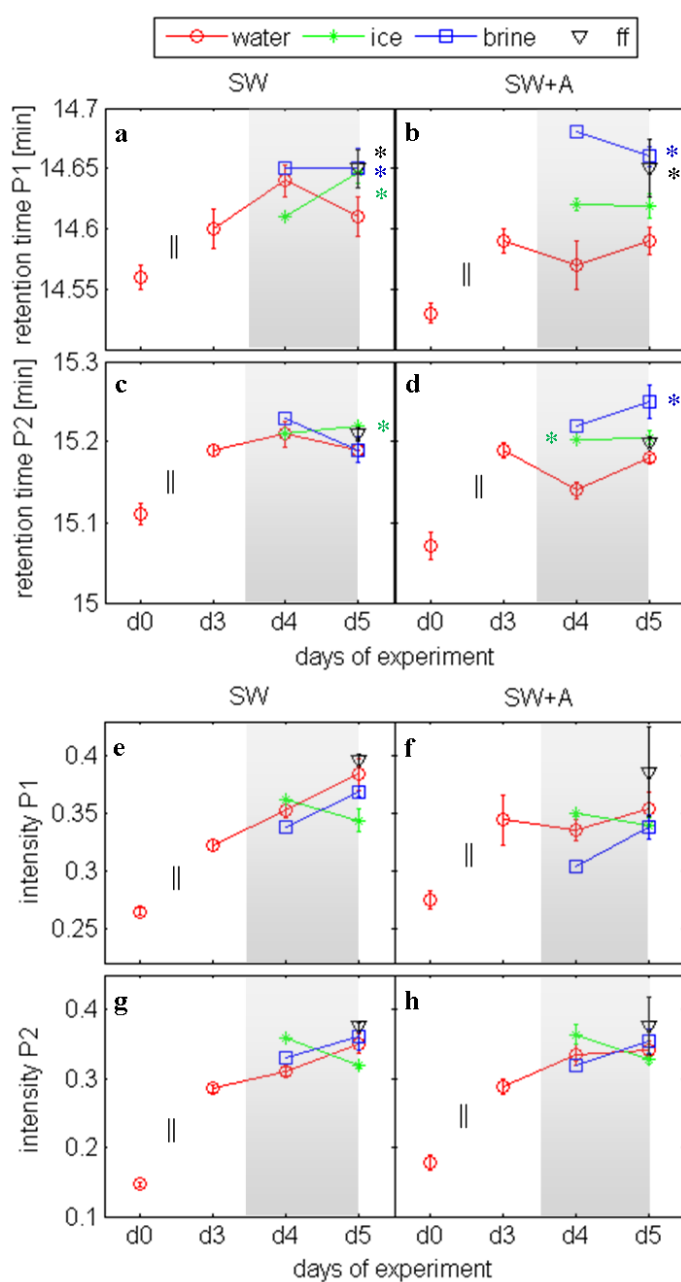


Fig. 10: The retention times (a-d) and absorbances (e-f) of peak 1 (P1) and 2 (P2) in the LC-SEC chromatograms. Full chromatograms are exemplified in Fig. 2 c. See Fig. 4 for more detailed explanation.

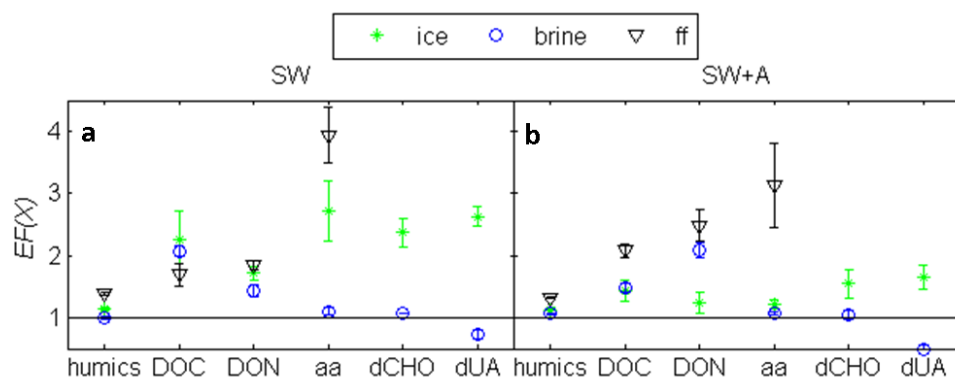


Fig.11: The enrichment factors (EF) of DOM fractions, humic like DOM, DOC, DON, amino acid-like fluorophores (aa), dissolved carbohydrates (dCHO) and dissolved uronic acids (dUA), in bulk ice, brine and frost flowers in each treatment (SW or SW+A) on d4 and d5. The black line shows the zero enrichment. The markers show the means and standard error among replicate mesocosms (n = 1 - 3).